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NEWS 1	Web Page for STN Seminar Schedule - N. America
NEWS 2	DEC 01 ChemPort single article sales feature unavailable
NEWS 3	JUN 01 CAS REGISTRY Source of Registration (SR) searching enhanced on STN
NEWS 4	JUN 26 NUTRACEUT and PHARMAML no longer updated
NEWS 5	JUN 29 IMSCOPROFILE now reloaded monthly
NEWS 6	JUN 29 EPFULL adds Simultaneous Left and Right Truncation (SLART) to AB, MCLM, and TI fields
NEWS 7	JUL 09 PATDPAFULL adds Simultaneous Left and Right Truncation (SLART) to AB, CLM, MCLM, and TI fields
NEWS 8	JUL 14 USGENE enhances coverage of patent sequence location (PSL) data
NEWS 9	JUL 27 CA/Caplus enhanced with new citing references
NEWS 10	JUL 16 GBFULL adds patent backfile data to 1855
NEWS 11	JUL 21 USGENE adds bibliographic and sequence information
NEWS 12	JUL 28 EPFULL adds first-page images and applicant-cited references
NEWS 13	JUL 28 INPADOCDB and INPAFAMDB add Russian legal status data
NEWS 14	AUG 08 Improve STN by completing a survey and be entered to win a gift card
NEWS 15	AUG 10 Time limit for inactive STN sessions doubles to 40 minutes

NEWS EXPRESS MAY 26 09 CURRENT WINDOWS VERSION IS V8.4,  
AND CURRENT DISCOVER FILE IS DATED 06 APRIL 2009.

NEWS HOURS STN Operating Hours Plus Help Desk Availability  
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\* Please take a couple of minutes to complete our short survey. Your  
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\* See NEWS 14 for details or go directly to the survey at:  
\* <http://www.zoomerang.com/Survey/?p=WEB229H4S9Q5UL>

FILE 'HOME' ENTERED AT 13:55:47 ON 14 AUG 2009

=> FIL REG  
COST IN U.S. DOLLARS  
SINCE FILE  
ENTRY SESSION  
0.22 0.22  
FULL ESTIMATED COST

FILE 'REGISTRY' ENTERED AT 13:56:13 ON 14 AUG 2009  
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STRUCTURE FILE UPDATES: 13 AUG 2009 HIGHEST RN 1174270-19-98  
DICTIONARY FILE UPDATES: 13 AUG 2009 HIGHEST RN 1174270-19-98

New CAS Information Use Policies, enter HELP USAGETERMS for details.

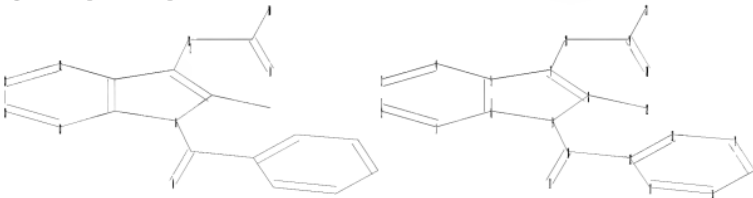
TSCA INFORMATION NOW CURRENT THROUGH June 26, 2009.

Please note that search-term pricing does apply when conducting SmartSELECT searches.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

<http://www.cas.org/support/stn/gen/stndoc/properties.html>

=> Uploading C:\Program Files\STNEXP\Queries\10541429\INDOELS.str

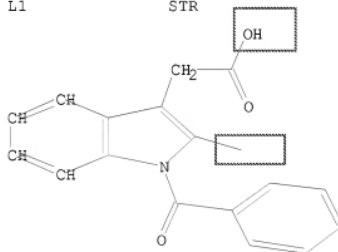


chain nodes :  
 10 11 18 19 20 21 22  
 ring nodes :  
 1 2 3 4 5 6 7 8 9 12 13 14 15 16 17  
 chain bonds :  
 7-18 8-22 9-10 10-11 10-12 18-19 19-20 19-21  
 ring bonds :  
 1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-9 7-8 8-9 12-13 12-17 13-14 14-15 15-16  
 16-17  
 exact/norm bonds :  
 5-7 6-9 7-8 8-9 9-10 10-11  
 exact bonds :  
 7-18 8-22 10-12 18-19  
 normalized bonds :  
 1-2 1-6 2-3 3-4 4-5 5-6 12-13 12-17 13-14 14-15 15-16 16-17 19-20  
 19-21

Match level :  
 1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:CLASS  
 11:CLASS 12:Atom 13:Atom 14:Atom 15:Atom 16:Atom 17:Atom 18:CLASS 19:CLASS  
 20:CLASS 21:CLASS 22:CLASS

L1        STRUCTURE UPLOADED

=> D  
 L1 HAS NO ANSWERS  
 L1                    STR



Structure attributes must be viewed using STN Express query preparation.

=> S L1 FULL  
 FULL SEARCH INITIATED 13:56:33 FILE 'REGISTRY'  
 FULL SCREEN SEARCH COMPLETED -        1754 TO ITERATE

100.0% PROCESSED        1754 ITERATIONS  
 SEARCH TIME: 00.00.01

123 ANSWERS

L2 123 SEA SSS FUL L1

=> FIL CAPLUS	SINCE FILE	TOTAL
COST IN U.S. DOLLARS	ENTRY	SESSION
FULL ESTIMATED COST	185.88	186.10

FILE 'CAPLUS' ENTERED AT 13:56:37 ON 14 AUG 2009  
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FILE COVERS 1907 - 14 Aug 2009 VOL 151 ISS 8  
FILE LAST UPDATED: 13 Aug 2009 (20090813/ED)  
REVISED CLASS FIELDS (NCL) LAST RELOADED: Jun 2009  
USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Jun 2009

Cplus now includes complete International Patent Classification (IPC) reclassification data for the second quarter of 2009.

CAS Information Use Policies apply and are available at:

<http://www.cas.org/legal/infopolicy.html>

This file contains CAS Registry Numbers for easy and accurate substance identification.

The ALL, BIB, MAX, and STD display formats in the CA/CAplus family of databases have been updated to include new citing references information. This enhancement may impact record import into database management software. For additional information, refer to NEWS 9.

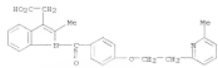
=> S L2
L3 29 L2
=> D IBIB 1-10



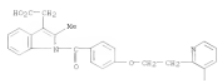




L3 ANSWER 7 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

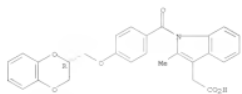


522404-23-1 CAPLUS  
1H-Indole-3-acetic acid, 2-methyl-1-(4-(2-(3-methyl-2-pyrrolidinylmethoxy)benzoyl)-2-methyl-1H-indole-3-acetic acid, (CA INDEX NAME)



522404-24-1 CAPLUS  
1H-Indole-3-acetic acid, 1-(4-((2S)-2,3-dihydro-1,4-benzodioxin-2-yl)methoxy)benzoyl-2-methyl-1H-indole-3-acetic acid, (CA INDEX NAME)

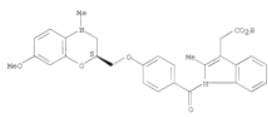
Absolute stereochemistry.



522404-25-1 CAPLUS  
1H-Indole-3-acetic acid, 1-(4-((2S)-2,3-dihydro-1,4-benzodioxin-2-yl)methoxy)benzoyl-2-methyl-1H-indole-3-acetic acid, (CA INDEX NAME)

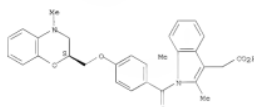
L3 ANSWER 7 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

Absolute stereochemistry.



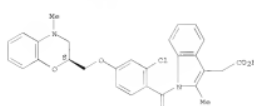
522404-54-4 CAPLUS  
1H-Indole-3-acetic acid, 1-(4-((2S)-3,4-dihydro-4-methyl-2H-1,4-benzodioxin-2-yl)methoxy)benzoyl-2-methyl-1H-indole-3-acetic acid, (CA INDEX NAME)

Absolute stereochemistry.

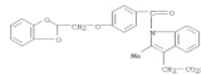


522404-59-9 CAPLUS  
1H-Indole-3-acetic acid, 1-(2-chloro-4-((2S)-3,4-dihydro-4-methyl-2H-1,4-benzodioxin-2-yl)methoxy)benzoyl-2-methyl-1H-indole-3-acetic acid, (CA INDEX NAME)

Absolute stereochemistry.

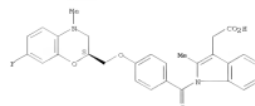


L3 ANSWER 7 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)



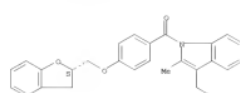
522404-64-6 CAPLUS  
1H-Indole-3-acetic acid, 1-(4-((2S)-3,4-dihydro-4-methyl-2H-1,4-benzodioxin-2-yl)methoxy)benzoyl-2-methyl-1H-indole-3-acetic acid, (CA INDEX NAME)

Absolute stereochemistry.



522404-67-7 CAPLUS  
1H-Indole-3-acetic acid, 1-(4-((2S)-3,4-dihydro-2H-1,4-benzodioxin-2-yl)methoxy)benzoyl-2-methyl-1H-indole-3-acetic acid, (CA INDEX NAME)

Absolute stereochemistry.

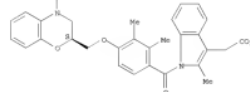


522404-68-8 CAPLUS  
1H-Indole-3-acetic acid, 1-(4-((2S)-3,4-dihydro-4-methyl-2H-1,4-benzodioxin-2-yl)methoxy)benzoyl-2-methyl-1H-indole-3-acetic acid, (CA INDEX NAME)

L3 ANSWER 7 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

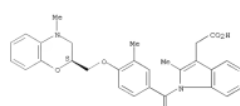
522404-69-3 CAPLUS  
1H-Indole-3-acetic acid, 1-(4-((2S)-3,4-dihydro-4-methyl-2H-1,4-benzodioxin-2-yl)methoxy)benzoyl-2-methyl-1H-indole-3-acetic acid, (CA INDEX NAME)

Absolute stereochemistry.



522404-69-3 CAPLUS  
1H-Indole-3-acetic acid, 1-(4-((2S)-3,4-dihydro-4-methyl-2H-1,4-benzodioxin-2-yl)methoxy)benzoyl-2-methyl-1H-indole-3-acetic acid, (CA INDEX NAME)

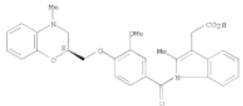
Absolute stereochemistry.



522404-63-5 CAPLUS  
1H-Indole-3-acetic acid, 1-(4-((2S)-3,4-dihydro-4-methyl-2H-1,4-benzodioxin-2-yl)methoxy)benzoyl-2-methyl-1H-indole-3-acetic acid, (CA INDEX NAME)

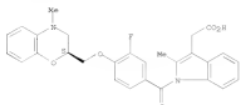
Absolute stereochemistry.

L3 ANSWER 7 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)



502404-54-5 CAPLUS  
1b-Indole-3-acetic acid, 1-[4-[(2b)-3,4-dihydro-4-methyl-2H-1,4-benzodiazin-2-yl)methoxy]-3-(furosemoyl)-2-methyl- (CA INDEX NAME)

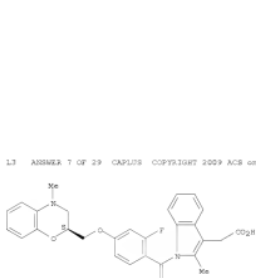
Absolute stereochemistry.



502404-55-7 CAPLUS  
1b-Indole-3-acetic acid, 1-[4-[(2b)-3,4-dihydro-4-methyl-2H-1,4-benzodiazin-2-yl)methoxy]-2-methoxybenzoyl]-2-methyl- (CA INDEX NAME)

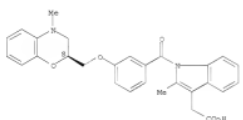
Absolute stereochemistry.

L3 ANSWER 7 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)



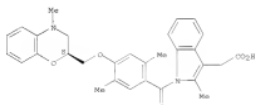
502404-56-9 CAPLUS  
1b-Indole-3-acetic acid, 1-[3-[(2b)-3,4-dihydro-4-methyl-2H-1,4-benzodiazin-2-yl)methoxy]benzoyl]-2-methyl- (CA INDEX NAME)

Absolute stereochemistry.



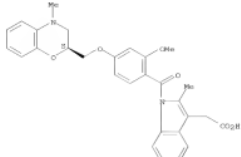
502404-62-3 CAPLUS  
1b-Indole-3-acetic acid, 1-[4-[(2b)-3,4-dihydro-4-methyl-2H-1,4-benzodiazin-2-yl)methoxy]-2,3-dimethylbenzoyl]-3-methyl- (CA INDEX NAME)

Absolute stereochemistry.



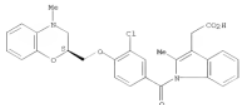
502404-84-0 CAPLUS  
1b-Indole-3-acetic acid, 3-[(2b)-3,4-dihydro-4-methyl-2H-1,4-benzodiazin-2-yl)methoxy]benzoyl]-2-methyl- (CA INDEX NAME)

L3 ANSWER 7 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)



502404-66-8 CAPLUS  
1b-Indole-3-acetic acid, 1-[3-chloro-4-[(2b)-3,4-dihydro-4-methyl-2H-1,4-benzodiazin-2-yl)methoxy]benzoyl]-2-methyl- (CA INDEX NAME)

Absolute stereochemistry.

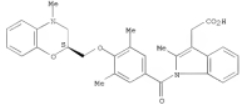


502404-67-9 CAPLUS  
1b-Indole-3-acetic acid, 1-[4-[(2b)-3,4-dihydro-4-methyl-2H-1,4-benzodiazin-2-yl)methoxy]-2-fluorobenzoyl]-2-methyl- (CA INDEX NAME)

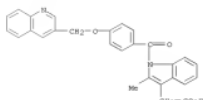
Absolute stereochemistry.

L3 ANSWER 7 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)  
benzodiazin-2-yl)methoxy]-7,5-dimethylbenzoyl]-2-methyl- (CA INDEX NAME)

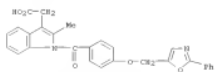
Absolute stereochemistry.



502407-09-2 CAPLUS  
1b-Indole-3-acetic acid, 2-methyl-1-[4-(3-quinolinylmethoxy)benzoyl]- (CA INDEX NAME)



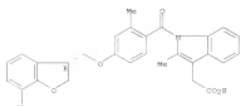
502407-11-6 CAPLUS  
1b-Indole-3-acetic acid, 2-methyl-1-[4-[(2-phenyl-5-oxazolyl)methoxy]benzoyl]- (CA INDEX NAME)



502407-13-8 CAPLUS  
1b-Indole-3-acetic acid, 2-methyl-1-[4-[(1-methyl-1H-indol-2-yl)methoxy]benzoyl]- (CA INDEX NAME)



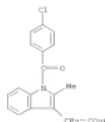
L3 ANSWER 7 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)



OS\_CITING REF COUNT: 8 THERE ARE 9 CAPLUS RECORDS THAT CITE THIS RECORD  
 (17 CITINGS)  
 REFERENCE COUNT: 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE REFORMAT

L3 ANSWER 8 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 20021903734 CAPLUS  
 DOCUMENT NUMBER: 16790-34-4  
 TITLE: Structure-activity relationship of substituted 2-methyl-3-oxo-2-azabicyclo[3.1.0]hex-5-ene-5-carboxylic acids and anti-inflammatory activity of 1-alkyl-3-methyl-5-prorrolones  
 AUTHOR(S): Department of Pharmaceutical Chemistry, Faculty of Pharmacy, New Delhi, 110 026, India  
 SOURCE: J. Indian Chem. Soc. (2002), 18(2), 375-376  
 PUBLISHER: Oriental Scientific Publishing Co.  
 PUBLICATION TYPE: Journal Article  
 LANGUAGE: English  
 OTHER INFO: (17 CITINGS) 159-28-76  
 AS\_CITING REF COUNT: 1-alkyl-3-methyl-5-prorrolones were prepared by converting the carbonylic acids to their hydrates and cyclizing them with MeCOCl/DMF. The anti-inflammatory activity of indomethacin in the rat paw edema test was nearly equal to that of indomethacin in the rat paw edema test.  
 IT 16790-34-4, 2-Methyl-1-(4-chlorobenzoyl)-3-indoleacetic acid  
 JAI 16790-34-4, 1-(4-chlorobenzoyl)-3-indoleacetic acid (Benzodiazepine derivative)  
 (preparation and anti-inflammatory activity of 1-alkyl-3-methyl-5-prorrolones)  
 PR 16790-34-4, 1-(4-chlorobenzoyl)-3-indoleacetic acid  
 CH 1B-Indole-3-acetic acid, 1-(4-chlorobenzoyl)-2-methyl- (CA INDEX NAME)



OS\_CITING REF COUNT: 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD  
 (1 CITINGS)  
 REFERENCE COUNT: 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE REFORMAT

L3 ANSWER 9 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 20021541130 CAPLUS

DOCUMENT NUMBER: 16791  
 TITLE: Structure-activity relationship of indomethacin analogues for MOP-1, COX-1 and COX-2 inhibition in human tumor cell line drug resistance modulators

AUTHOR(S): TOLMACH, J. L.; KARAS, M. M.; KARAS, A. I.; COHEN, M. J.; KARAS, R. J.; KARAS, A. I.; Dublin City University, The National Cell and Tissue Culture Centre, Glasnevin, Dublin 11, Ireland  
 SOURCE: J. Indian Chem. Soc. (2002), 18(2), 1651-1670  
 PUBLISHER: Indian Chemical Society  
 PUBLICATION TYPE: Journal Article  
 LANGUAGE: English  
 ABSTRACT: The authors report the screening of analogs of indomethacin to investigate the structure-activity relationship (SAR) of indomethacin-mediated drug resistance associated protein-1 (MRP-1) inhibition. By examining the SAR of compounds with minor variations of the parent structure, the authors were able to sep. MRP-1, glutathione-S-transferase (GST), cyclooxygenase (COX)-1 and COX-2 inhibitory activities. Combination studies were performed to determine the additive or synergistic anti-inflammatory potential in MOP-1-expressing cell lines. MOP-1 (inside Out Vesicles (IOVs) were utilized to demonstrate the ability of the compounds to inhibit MRP-1. Most of the indomethacin analogs active as MRP-1 inhibitors were poor GST inhibitors when compared with the anti-inflammatory activity of indomethacin. Two of the indomethacin analogs were found to have high COX-1 inhibitory activity and low COX-2 inhibitory activity, suggesting potentially reduced toxicity. One MRP-1 inhibitory indomethacin analog was also found to have low COX-1 inhibitory activity, but significant COX-2 inhibitory activity. This analog was found to have reduced MRP-1 mediated cellular toxicity, but with the possibility of direct inhibitory effects on tumor growth.

IT 16791-34-4  
 ELI PAC (Pharmacological activity); PRP (Properties); THER (Therapeutic use); (17 CITINGS)  
 TITLE: Structure-activity relationship of indomethacin analogs for MOP-1, GST, COX-1 and COX-2 inhibition identification of novel drug resistance modulators in human tumor cell line  
 PR 16791-34-4, CAPLUS  
 CH 1B-Indole-3-acetic acid, 1-(4-chlorobenzoyl)-2-methyl- (CA INDEX NAME)

L3 ANSWER 9 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

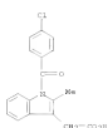
compound 25 in pub.

OS\_CITING REF COUNT: 18 THERE ARE 18 CAPLUS RECORDS THAT CITE THIS RECORD (18 CITINGS)  
 REFERENCE COUNT: 35 THERE ARE 35 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE REFORMAT



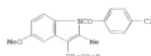
L3 ANSWER 13 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)  
 130  $\mu\text{M}$  for a COX-2 selectivity index of 8.4. Binding to the center of the COX-2 active site of human COX-2 showed it binds in the center of the COX-2 binding site with the C-5 (Ph) ring oriented toward the acetylation site (Ser530), the Ph group of the C-4 (benzyl) moiety oriented in the vicinity of the COX-2 secondary binding pocket near Val153.

131 16390-26-4  
 132 AIV (Adverse effect, including toxicity); TEF (Therapeutic use); ECOL (Ecological/Environmental); SAR of aranacyclopeptins, novel COX-2 inhibitors; antiinflammatories  
 133 16390-26-4 CAS#  
 134 18-Indole-3-acetic acid, 1-(4-chlorobenzoyl)-2-methyl- (CA INDEX NAME)



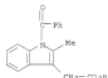
CS\_CITING KEY COUNT: 5 THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD  
 REFERENCE COUNT: 39 THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE REFORMAT

L3 ANSWER 13 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN  
 135 ACCESSION NUMBER: 1983141682 CARLOS  
 DOCUMENT NUMBER: 99134670  
 ORIGINAL REFERENCE NO.: 93138273a,182904  
 TITLE: Pharmacokinetic studies of delmetethacin and indometethacin in rats  
 AUTHOR(S): Bdg, Inst. Farmacocinet. Metab., J. Uriach y Cia, S. A., Barcelona, Spain  
 SOURCE: Rev. de Farmacología y Toxicología (1982), 8(2), 261-4  
 DOCUMENT TYPE: Article  
 LANGUAGE: Spanish  
 C2

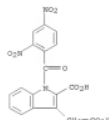


136 The pharmacokinetics of indometethacin (I) [57-86-1] and delmetethacin (II) [16390-26-1] were compared in rat studies, with both drugs given i.v. and orally. Both drugs were rapidly absorbed and eliminated, with I and II exhibiting noncompartmental kinetics. The rapid elimination of II and the lack of deep compartments imply a low tendency to form deposits or reservoirs which could result in toxic effects. I, however, exhibited triponential kinetics and a longer half-life, so the risk of accumulation was greater. Delmetethacin (II) is a prodrug of I. Delmetethacin compared with I is consistent with the different pharmacokinetic behavior of the 2 compounds. Delmetethacin is a prodrug of I, which has easier access to those sites where its anti-inflammatory activity is needed.  
 137 16401-90-2  
 138 Hs (Biological process); BSY (Biological study, unclassified); ECOL (Biological study); PROC (Process)  
 139 16401-90-2 CAS#  
 140 18-Indole-3-acetic acid, 1-benzoyl-2-methyl- (CA INDEX NAME)

L3 ANSWER 13 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)



L3 ANSWER 14 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN  
 141 ACCESSION NUMBER: 1980151424 CARLOS  
 DOCUMENT NUMBER: 99138277a,182904  
 ORIGINAL REFERENCE NO.: 93138277a,182904  
 TITLE: A convenient synthesis of new indole derivatives  
 AUTHOR(S): S. A. Khan, M. S. Siddiqui, Amin A., Khan, Nasreen B., Dep. Chem., Aligarh Muslim Univ., Aligarh, 202 001, India  
 SOURCE: Indian Journal of Chemistry, Section B: Organic Chemistry Including Medicinal Chemistry (1980), 19B, 125-8  
 DOCUMENT TYPE: Journal  
 LANGUAGE: English  
 AB Several 1-substituted indole derivs. have been prepared by refluxing equimolar amt. of the appropriate indole and 2,4-dinitrobenzoyl chloroformate in the presence of NaOAc.  
 17 74693-46-2  
 18 E1 (Synthetic preparation); PREP (Preparation)  
 19 74693-46-2 (preparation of)  
 20 74693-46-2 CAS#  
 21 18-Indole-3-acetic acid, 2-carboxy-1-(2,4-dinitrobenzoyl)- (CA INDEX NAME)



CS\_CITING KEY COUNT: 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD  
 (1 CITINGS)

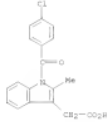




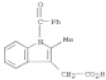
1.1 ANSWER 19 OF 29 CAPTION COPYRIGHT 2009 ACS ON 27N (Continued)  
JP 1966-3187 A 1966012  
NO 1966-162587 A 1966041  
PT 1966-945 A 1966041

test in rats II had an oral ED50 of 80 mg/kg and a therapeutic ratio of 18.5. IT 16400-26-4P 16401-80-2P 16401-91-3P  
16401-83-5P NL: SM (Synthetic preparation); PMP (Preparation)

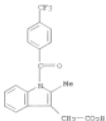
328 (preparation of)  
16390-26-6 CAPTUS  
CB 16-Indole-1-acetic acid, 1-(4-chlorobenzoyl)-2-methyl- (CA INDEX NAME)



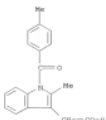
322 16401-80-2 CAPLOS  
C21 18-Indole-3-acetic acid, 1-benzoyl-2-methyl- ICA INDEX NAME



INN 164-01-81-3 CAPLOSS  
CN 1B-Indole-3-acetic acid, 2-methyl-1-[4-(trifluoromethyl)benzoyl]-  
INDOLE NAME) (CA)



BN 16401-83-5 CAPLOS  
CN 1H-Indole-3-acetic acid, 2-methyl-1-(4-methylbenzoyl)- (CA INDEX NAME)



05.CITING REF COUNT: 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD

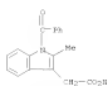
13 ANSWER 19 OF 29 CAPTION COPYRIGHT 2009 ACS on 8/7/09 (Continued)  
[1 CITATIONS]

LC ANSWER 20 OF 29 CAPUS COPIRIGHT 2009 ACS ON 87W  
ACCESSION NUMBER 197445943 CAPUS  
ORIGINAL REFERENCE NO. 0179124, 7914a  
TITLE: *N*-Methoxycarbonyl phenylhydrazines  
from *Phenylhydrazine*, Maru  
PATENT ASSIGNEE(S): Suntoku Chemical Co., Ltd.  
SOURCE: U.S., 12 pp. Division of U.S. 3,629,284 (CA  
W/112040G).  
COUNTRY: KOREAN  
DOCUMENT TYPE: LANGUAGES: English  
PCT ACC. NO.: COUNTRY: ?  
PATENT INFORMATION:  
PATENT INFORMATION:

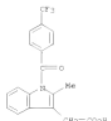
GI For diagram(s), see printed CA issues.  
 AB  $\text{ROOC}(\text{NH}_2)\text{C}_6\text{H}_4\text{R}$  [I, R =  $p$ -C<sub>6</sub>H<sub>4</sub>,  $p$ -MeC<sub>6</sub>H<sub>4</sub>, Ph,  $p$ -MeOC<sub>6</sub>H<sub>4</sub>,  $p$ -F<sub>3</sub>C<sub>6</sub>H<sub>4</sub>,  $p$ -BrC<sub>6</sub>H<sub>4</sub>,  $p$ -FC<sub>6</sub>H<sub>4</sub>, 3-pyridyl, 4-pyridyl, 2-thienyl, 5-chloro-2-thienyl, 2-furyl,  $p$ -MeOC<sub>6</sub>H<sub>4</sub>, 2-naphthyl; R' = H,  $p$ -Cl,  $p$ -Me,  $p$ -MeO,  $p$ -F,  $p$ -Me,  $p$ -MeO,  $p$ -F,  $p$ -Me]



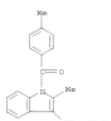
L3 ANSWER 21 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)  
 CS 18-Indole-3-acetic acid, 1-benzoyl-3-methyl- (CA INDEX NAME)



NR 16401-81-3 CAPLUS  
 CS 18-Indole-3-acetic acid, 2-methyl-1-[4-(trifluoromethyl)benzoyl]- (CA INDEX NAME)



NR 16442-92-5 CAPLUS  
 CS 18-Indole-3-acetic acid, 2-methyl-1-(4-methoxybenzoyl)- (CA INDEX NAME)



L3 ANSWER 22 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 19711215421 CAPLUS

DOCUMENT TYPE: JP 45037528

ORIGINAL REFERENCE NO.: 74120229a,20242a

TITLE: 1-Arylindole derivatives

INVENTOR(S): Tsuru, Toshiaki; Nakao, Masaru

PATENT ASSIGNEE(S): Sumitomo Chemical Co., Ltd.

SCOPC: Jpn. Tokkyo Koho, 3 pp.

DOCUMENT TYPE: Patents

LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 45037528	JP	39703128	JP	19470512

GI For diazepam, see printed CA Issue.

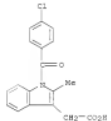
AB 1, useful as an antiinflammatory, analgesic, and antipyretic, is prepared by a simple method. For example, 1-(4-chlorobenzoyl)-3-methylindole and acetosuccinic acid in *H*OH are warmed 4 hr at 95-96° to give 1-(4-chlorobenzoyl)-3-methyl-1,2-dihydro-2-methyl-1,2-dihydro-3-indoleacetic acid (MeCO<sub>2</sub>, R2 = Me), m. 150-151° (aqueous MeCO<sub>2</sub>). Similarly prepared are 9 adducts.

IT 16390-26-42

ZK 16390-26-42 (Synthetic preparation); PAPD (Preparation)

[preparation of]

NR 16390-26-4 CAPLUS  
 CS 18-Indole-3-acetic acid, 1-(4-chlorobenzoyl)-2-methyl- (CA INDEX NAME)



L3 ANSWER 23 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)  
 CS\_CITING REF COUNT: 4 THERE ARE 4 CAPLUS RECORDS THAT CITE THIS RECORD  
 (4 CITATIONS)



L3 ANSWER 23 OF 29 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 19711215722 CAPLUS

DOCUMENT TYPE: JP 45037522

ORIGINAL REFERENCE NO.: 74124245a,14252a

TITLE: 1-Arylindoleacetic acid derivatives

INVENTOR(S): Wakimura, Atsushi; Nakamura, Yasushi; Nakao, Masaru

PATENT ASSIGNEE(S): Sumitomo Chemical Co., Ltd.

SCOPC: JP 45037522 (Synthetic) 4 pp.

DOCUMENT TYPE: Patents

LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 45037522	JP	39703128	JP	19470522

GI For diazepam, see printed CA Issue.

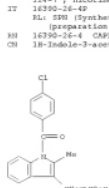
AB 1, useful as an antiinflammatory, analgesic, and antipyretic, are manufactured. For example, 1-(4-chlorobenzoyl)-3-methylindole and acetosuccinic acid in *H*OH are warmed 4 hr at 95-96° to give 1-(4-chlorobenzoyl)-3-methyl-1,2-dihydro-2-methyl-1,2-dihydro-3-indoleacetic acid (MeCO<sub>2</sub>, R2 = Me), m. 150-151° (aqueous MeCO<sub>2</sub>). Similarly prepared are 9 adducts.

IT 16390-26-42

ZK 16390-26-42 (Synthetic preparation); PAPD (Preparation)

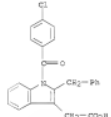
[preparation of]

NR 16390-26-4 CAPLUS  
 CS 18-Indole-3-acetic acid, 1-(4-chlorobenzoyl)-2-methyl- (CA INDEX NAME)









OS.CITING RLF COUNT: 5 THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD

L3 ANSWER 23 OF 23 CAPTION: COPRIGHT 2009 ACS OR STM  
ACCESSION NUMBER: 1974-690668 CAPLOS  
PUBLICATION NUMBER: 66  
ORIGINAL PREFERENCE NO.: 6717053a,17062a  
TITLE: Preparation of 1-acyl-3-alkoxy carbonyl alkylidene malonate monobiles  
PATENT ASSIGNEE(S): Suntomo Chemical Co., Ltd.  
SOURCE: Chempat NAKAJI  
DOCUMENT TYPE: Patent  
LANGUAGE: Dutch  
PUBLICATION NUMBER: 66

L3 ANSWER 29 OF 29 CARLOS COPYRIGHT 2009 ACS on STN (Continued)

JP 1945-81735	A 19651229
JP 1945-81796	A 19651229
JP 1946-3187	A 19660120
JP 1946-5754	A 19660131
JP 1946-7276	A 19660207
JP 1946-7277	A 19660207
NO 1946-16287	A 19660414
PT 1946-995	A 19660418
NL 1946-5169	AT 19660418

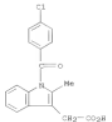
CC For diagram(s), see printed OA page.  
 At 7b - *isobutyl*, Py - *pyridyl*, Ph - *aryl*, and d - decomposition throughout this abstract. The title compound, (1) are antiinflammatory, antiipyretic and analgesic compounds which are prepared by the reaction of N-acetylated phenylhydrazine (11) with an oxo acid  $\text{ACOCOC}(\text{CH}_2\text{N}(\text{CH}_3)\text{CO})\text{CO}^+$ .

II is obtained by decomposition of hydrazones (III), which is obtained by acylation of IV with ARCON (X is halogen or ester residue). Thus, to a solution of 1 g. IV (X = *p*-MeO, X = H, X = *Me*) in 30 ml. pyridine, 15 g. 4-Chloro-*tert*-butyl-*tert*-butylcarboxylic acid is added dropwise with ice cooling. The reaction mixture is heated at room temperature and poured into ice-H<sub>2</sub>O to give 19 g. III (X = *p*-MeO, X = *Me*, Ar = *p*-ClC<sub>6</sub>H<sub>4</sub>), m. 157-8° (EtOH, H<sub>2</sub>O). To a solution of 3.4 g. IV (X = *p*-MeO, X = H, X = *Me*) in 10 ml. pyridine, 10 g. 4-Chloro-4*tert*-butyl-4*tert*-butylcarboxylic acid is added with ice-cooling. The mixture is left at room temperature and concentrated into ice-H<sub>2</sub>O to give 2.5 g. III (X = *p*-MeO, Ar = *p*-ClC<sub>6</sub>H<sub>4</sub>), m. 157-2°. A solution of 9.5 g. V in 90 ml. EtOH is saturated with HCl. The mixture is left at ambient temperature, concentrated, and worked up to give VI. A solution of 9.5 g. VI and 11.4 g. levulinic acid is heated 3 hrs. at 75° (DAGMAR, CHT-100). The mixture is left at ambient temperature and poured into EtOH to give 14.5 g. VII (X = *p*-Py, X = *Me*, X = *p*-MeO, n = *p* = *Q*, n = 1, *R* = OH) (VII), n. 181-9° (Me<sub>2</sub>CO, H<sub>2</sub>O) (method a). In method b AcOH is used as the solvent. A mixture of 9 g. VI, 4.2 g. Me levulinate, and 40 ml. MeOH is refluxed 5 hrs. with stirring. The MeOH is evaporated in vacuo and the precipitate worked up to give VII Me ester (VII), n. 113-15° (MeOH) (method a). A mixture of 1 g. VII-HCl (Ar = *p*-ClC<sub>6</sub>H<sub>4</sub>, X = *p*-MeO) and 1 g.

L3 ANSWER 29 OF 29 CARLOS COPYRIGHT 2009 ACS on STN (Continued)

gives 1.4 g. VII (X = <i>p</i> -MeO, Ar = <i>p</i> -ClC <sub>6</sub> H <sub>4</sub> ), m. 113-15° (MeOH) (method b). The product is dried and the residue worked up to give VIII (method c). III (X = <i>p</i> -MeO, Ar = <i>p</i> -ClC <sub>6</sub> H <sub>4</sub> , R = <i>Me</i> ) is obtained by the reaction of 19 g. III (X = <i>p</i> -MeO, Ar = <i>p</i> -ClC <sub>6</sub> H <sub>4</sub> , R = <i>Me</i> ) and 4 g. dry HCl gas is passed with ice-cooling. The melt is heated slowly and refluxed 1.5 hrs. The EtOH is added to give a resin, which is dissolved in EtOH and filtered. The solution is concentrated in vacuo to give 10 g. III (X = <i>p</i> -MeO, Ar = <i>p</i> -ClC <sub>6</sub> H <sub>4</sub> , R = <i>Me</i> ), m. 143-145° (EtOH) (method d). The product is dried and the residue worked up to give VIII (method g). [TABLE OMITTED]
17 16399-26-4F 16401-90-2P 16401-91-3P
ELI 39M (Synthetic preparation); FEPF (Preparation) (preparation of)
CH 16399-14-4 CAPUS

CH 18-Indole-3-acetic acid, 1-(4-chlorobenzoyl)-2-methyl- (CA INDEX NAME)



CH 16401-90-2 CAPUS

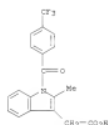
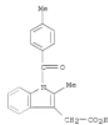
CH 18-Indole-3-acetic acid, 1-benzoyl-2-methyl- (CA INDEX NAME)

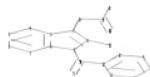
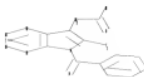


CH 16401-91-3 CAPUS

CH 18-Indole-3-acetic acid, 2-methyl-3-(4-(trifluoromethyl)benzoyl)- (CA INDEX NAME)

L3 ANSWER 29 OF 29 CARLOS COPYRIGHT 2009 ACS on STN (Continued)

CH 16401-83-5 CAPUS  
CH 18-Indole-3-acetic acid, 2-methyl-1-(4-methoxybenzoyl)- (CA INDEX NAME)



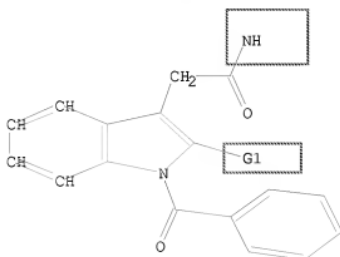
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 10 11 18 19 20 21 22  
 ring nodes :  
 1 2 3 4 5 6 7 8 9 12 13 14 15 16 17  
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 7-18 8-22 9-10 10-11 10-12 18-19 19-20 19-21  
 ring bonds :  
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 16-17  
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 normalized bonds :  
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G1:X,CH3,Et,n-Pr,i-Pr,n-Bu,i-Bu,s-Bu,t-Bu,CF3,CC12,CC13,CBr3

Match level :  
 1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:CLASS  
 11:CLASS 12:Atom 13:Atom 14:Atom 15:Atom 16:Atom 17:Atom 18:CLASS 19:CLASS  
 20:CLASS 21:CLASS 22:CLASS

L4 STRUCTURE UPLOADED

=> D  
 L4 HAS NO ANSWERS  
 L4 STR



G1 X, Me, Et, n-Pr, i-Pr, n-Bu, i-Bu, s-Bu, t-Bu, CF3, CC12, OC13, CBr3

Structure attributes must be viewed using STN Express query preparation.

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 SAMPLE SCREEN SEARCH COMPLETED - 60 TO ITERATE

100.0% PROCESSED 60 ITERATIONS 2 ANSWERS  
 SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*  
 BATCH \*\*COMPLETE\*\*  
 PROJECTED ITERATIONS: 736 TO 1664  
 PROJECTED ANSWERS: 2 TO 124

L5 2 SEA SSS SAM L4

=> S L4 FULL  
 FULL SEARCH INITIATED 14:15:51 FILE 'REGISTRY'  
 FULL SCREEN SEARCH COMPLETED - 1133 TO ITERATE

100.0% PROCESSED 1133 ITERATIONS 10 ANSWERS  
 SEARCH TIME: 00.00.01

L6 10 SEA SSS FUL L4

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DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
CA SUBSCRIBER PRICE	ENTRY	SESSION
	0.00	-20.50

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 USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Jun 2009

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=> S L6  
 L7                  4 L6  
 => D IBIB ABS HITSTR TOT

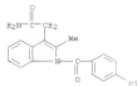




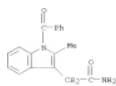
LT ANDERER 4 OF 4 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)  
 A-C(=O)(C)I-T heterocyclic A is a 5 to 8 membered heterocyclic ring where heterocyclic A is selected from N, O, S, etc.; the 5-6 membered heterocyclics ring is optionally fused with a second ring; T is NH2, NH-(cyclo)alkyl,  
 etc.; NH-cycloalkenyl, etc.), useful in inhibiting chemotactic activation of neutrophils and monocytes, and in the treatment of inflammatory diseases (IL-8) with CXCR1 and CXCR2 membrane receptors. The compds. are used for the prevention and treatment of pathologies deriving from said active principles.  
 In particular,  $\alpha$ -substituted arylacetic acids deriv., such as anilides and amides, etc., lack cyclo-olefinase inhibition activity and are particularly useful in the treatment of various types of pathologies such as psoriasis, ulcerative colitis, or melanoma, etc. For instance, prept. in the example 2 arylacetic acid deriv. 1 (10-09) showed 82% (IL-8) mediated inhibition activity on CXCR1 and CXCR2 receptors.  
 IT 740239-51-0 CAPLUS  
 ELI PAC (Pharmacological activity); SII (Synthetic preparation); TEC (Therapeutic use); ECOL (Biological study); PREP (Preparation); USES (Use); HMD (Heterocyclic compounds); HSC (Heterocyclic substances);  
 preparation of arylacetic acids useful for the treatment of IL-8  
 mediated diseases  
 22 740239-56-1 CAPLUS  
 23 1H-Indole-3-acetamide, 1-benzoyl-3-[3-(dimethylamino)propyl]-2-methyl- (CA INDEX NAME)



22 740239-56-1 CAPLUS  
 23 1H-Indole-3-acetamide, 1-benzoyl-3-[3-(dimethylamino)propyl]-2-methyl- (CA INDEX NAME)



LT ANDERER 4 OF 4 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)  
 NH 740239-57-0 CAPLUS  
 CH 1H-Indole-3-acetamide, 1-benzoyl-2-methyl- (CA INDEX NAME)



OC CITING REF COUNT: 2 THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD  
 REFERENCE COUNT: 3 (2 CITING)  
 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE REFORMAT